

10/739208

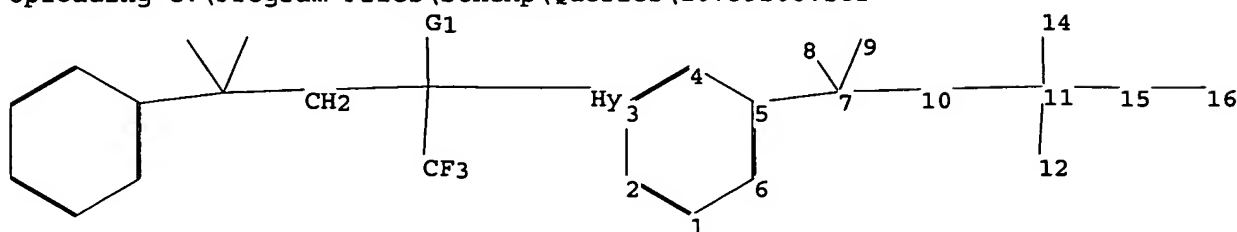
* * * * * * * * * * * * * * * STN Columbus * * * * * * * * * * * * * * *

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=> file reg

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Uploading C:\Program Files\Stnexp\Queries\10739208.str



chain nodes :

7 8 9 10 11 12 14 15 16

ring nodes :

1 2 3 4 5 6

chain bonds :

5-7 7-8 7-9 7-10 10-11 11-12 11-14 11-15 15-16

ring bonds :

1-2 1-6 2-3 3-4 4-5 5-6

exact/norm bonds :

11-14 15-16

exact bonds :

5-7 7-8 7-9 7-10 10-11 11-12 11-15

normalized bonds :

1-2 1-6 2-3 3-4 4-5 5-6

isolated ring systems :

containing 1 :

G1:OH,N

Match level :

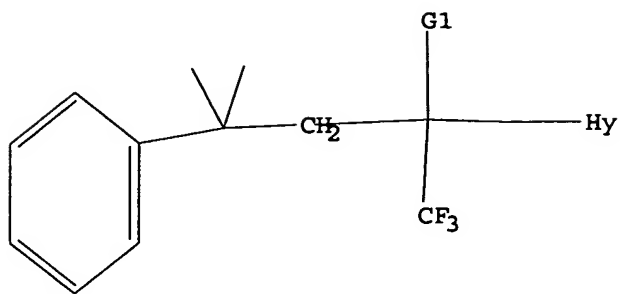
1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:CLASS 8:CLASS 9:CLASS 10:CLASS
11:CLASS 12:CLASS 14:CLASS 15:CLASS 16:Atom

L1 STRUCTURE UPLOADED

=> d l1

L1 HAS NO ANSWERS

L1 STR



G1 OH,N

Structure attributes must be viewed using STN Express query preparation.

=> s l1 full
L3 629 SEA SSS FUL L1

=> file ca

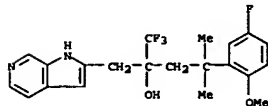
=> s l3
L4 7 L3

=> d ibib abs fhitstr 1-7

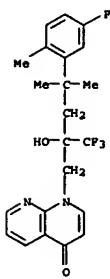
L4 ANSWER 1 OF 7 CA COPYRIGHT 2006 ACS on STN
 ACCESSION NUMBER: 144:184662 CA
 TITLE: Anti-aromatase compounds, pharmaceutical compositions, and use in the treatment of estrogen receptor-mediated disorders, including breast cancer and other cancers
 INVENTOR(S): Nelson, Richard More; Liu, Pingrong; Proudfoot, John Robert; Riether, Doris; Harcken, Christian Hanke Justus Joachim; Thomson, David S.
 PATENT ASSIGNEE(S): Boehringer Ingelheim Pharmaceuticals, Inc., USA
 SOURCE: U.S. Pat. Appl. Publ., 43 pp.
 CONDS: USXXCO
 DOCUMENT TYPE: Patent
 LANGUAGE: English
 FAMILY ACC. NUM. COUNT: 1
 PATENT INFORMATION:

| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|------------------------|------|----------|-----------------|------------|
| US 2006030608 | A1 | 20060209 | US 2005-137281 | 20050525 |
| PRIORITY APPLN. INFO.: | | | US 2004-598612P | P 20040804 |

OTHER SOURCE(S): MARPAT 144:184662
 AB The invention discloses aryl and heteroaryl alc. compds. (Markush included), or a tautomer, prodrug, solvate, or salt thereof, pharmaceutical compns. containing such compds., and methods for modulating estrogen receptor activity in a cell or patient or treating an estrogen receptor-mediated disorder, particularly breast and other cancers, in a patient in need thereof by administering an effective amount of compound of the invention.
 IT 609850-97-7
 RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses) (anti-aromatase compds., pharmaceutical compns., and use in treatment of estrogen receptor-mediated disorders)
 RN: 609850-97-7 CA
 CN: 1H-Pyrazolo[2,3-c]pyridine-2-ethanol, α -(2-(5-fluoro-2-methoxyphenyl)-2-methylpropyl)- α -(trifluoromethyl)- (9CI) (CA INDEX NAME)



L4 ANSWER 2 OF 7 CA COPYRIGHT 2006 ACS on STN (Continued)
 ylpentan-2-one is converted to the corresponding oxirane (DMSO, Me3SOI, NaH). This intermediate is oxidized to the methanesulfonyl analog and finally reacted with thieno[3,2-b]pyridin-7-ol (EtOH, NaOEt) to give II. Selected compds. of the invention exhibit potent activity in the glucocorticoid receptor binding assay. I are useful for the treatment of diseases and cardiovascular diseases.
 IT 866112-70-1P, 1-[4-(5-Fluoro-2-methylphenyl)-2-hydroxy-4-methyl-2-trifluoromethylpentyl]-1H-[1,8]naphthyridin-4-one
 RL: PAC (Pharmacological activity); RCT (Reactant); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); RACT (Reactant or reagent); USES (Uses) (preparation of naphthyridine-derived α -trifluoromethyl alcs. or amine and analogs as glucocorticoid mimetics)
 RN: 866112-70-1 CA
 CN: 1,8-Naphthyridin-4(iH)-one, 1-[4-(5-fluoro-2-methylphenyl)-2-hydroxy-4-methyl-2-(trifluoromethyl)pentyl]- (9CI) (CA INDEX NAME)

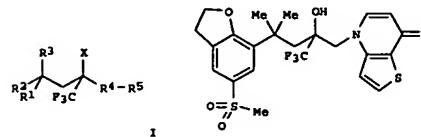


REFERENCE COUNT: 9 THERE ARE 9 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 3 OF 7 CA COPYRIGHT 2006 ACS on STN
 ACCESSION NUMBER: 143:367290 CA
 TITLE: Preparation of α -trifluoromethyl alcohols or amines as glucocorticoid mimetics
 INVENTOR(S): Regan, John Robinson; Lee, Thomas Mai-Ho; Thomson, David; Kirrane, Thomas Martin; Kuzmich, Daniel; Proudfoot, John Robert; Bekkali, Youness; Zindell, Renee
 PATENT ASSIGNEE(S): Boehringer Ingelheim Pharmaceuticals, Inc., USA
 SOURCE: PCT Int. Appl., 146 pp.
 DOCUMENT TYPE: Patent
 LANGUAGE: English
 FAMILY ACC. NUM. COUNT: 1
 PATENT INFORMATION:

| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|--|--|-----------|-----------------|------------|
| WO 2005095401 | A1 | 20050103 | WO 2005-US6975 | 20050104 |
| W: AB, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EO, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KB, KO, KP, KR, KW, LC, LK, LR, LS, LT, LV, MA, MD, MO, MK, MN, MM, MX, MZ, NA, NI, NO, NZ, OM, PG, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SM, SY, TJ, TM, TN, TR, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, | | | | |
| EW: | RW: BW, GH, GM, KE, LS, MM, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, PR, GB, GR, HU, IE, IS, IT, LT, LU, MC, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CO, CI, CM, GA, GN, GQ, GM, ML, MR, NE, SN, TD, TO | | | |
| US 2005214091 | A1 | 200501020 | US 2005-72819 | 20050304 |
| PRIORITY APPLN. INFO.: | | | US 2004-555220P | P 20040322 |

OTHER SOURCE(S): MARPAT 143:367290
 GI

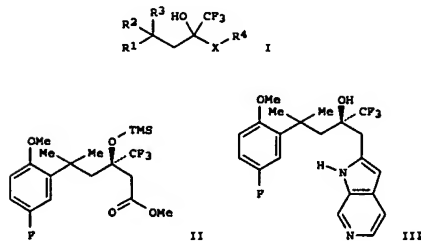


AB Title compds. I [R1 = (hetero)aryl, cycloalkyl, etc.; R2-3 = H, alkyl, arylalkyl, etc.; R4 = CO, divalent alkyl; R5 = 5-7 membered heterocyclic ring fused to a 5-7 membered heteroaryl/heterocyclic ring with one exception; X = OH, (un)substituted amino] are prepared. For instance, 1,1,1-trifluoro-4-methyl-4-(5-methylsulfanyl)-2,3-dihydrobenzofuran-7-

L4 ANSWER 3 OF 7 CA COPYRIGHT 2006 ACS on STN
 ACCESSION NUMBER: 143:325974 CA
 TITLE: Stereoselective synthesis of certain trifluoromethyl-substituted alcohols
 INVENTOR(S): Song, Jinhua J.; Tan, Zhulin; Yee, Nathan K.; Senanayake, Chris Hugh; Xu, Jinghua; Gallo, Fabrice
 PATENT ASSIGNEE(S): Boehringer Ingelheim Pharmaceuticals, Inc., USA
 SOURCE: U.S. Pat. Appl. Publ., 11 pp.
 DOCUMENT TYPE: Patent
 LANGUAGE: English
 FAMILY ACC. NUM. COUNT: 1
 PATENT INFORMATION:

| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|--|--|----------|-----------------|------------|
| US 200509488 | A1 | 20050922 | US 2005-70462 | 20050302 |
| WO 2005090343 | A1 | 20050929 | WO 2005-US6998 | 20050304 |
| W: AB, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KB, KO, KP, KR, KW, LC, LK, LR, LS, LT, LV, MA, MD, MO, MK, MN, MM, MX, MZ, NA, NI, NO, NZ, OM, PG, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SM, SY, TJ, TM, TN, TR, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, | | | | |
| ZW: | RW: BW, GH, GM, KE, LS, MM, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, PR, GB, GR, HU, IE, IS, IT, LT, LU, MC, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CO, CI, CM, GA, GN, GQ, GM, ML, MR, NE, SN, TD, TO | | | |
| PRIORITY APPLN. INFO.: | | | US 2004-554266P | P 20040318 |

OTHER SOURCE(S): CASREACT 143:325974; MARPAT 143:325974
 GI



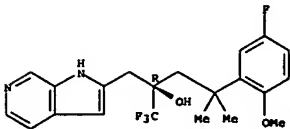
AB A process for stereoselective synthesis of I [R1 = (un)substituted aryl or heteroaryl; R2 and R3 = H or alkyl, or together from a spirocycle ring; X = (un)substituted alkyl, alkenyl, or alkynyl; R4 = (un)substituted

L4 ANSWER 3 OF 7 CA COPYRIGHT 2006 ACS on STN (Continued)
 heteroaryl group employing a chiral indane to control stereoselectivity with a novel ester to azaindole reaction in the last step. For example, the ester II (prepn. given) was reacted with 3-amino-4-picoline to provide the chiral alc. III in the direct ester to azaindole reaction step.

IT 865200-60-8
 RL: SPN (Synthetic preparation); PRSP (Preparation)
 (stereoselective prep of trifluoromethyl-substituted alcs. employing a chiral indane reactant)

RN 865200-60-8 CA
 CN 1H-Pyrido[2,3-c]pyridine-2-ethanol, α -(2-(5-fluoro-2-methoxyphenyl)-2-methylpropyl)- α -(trifluoromethyl)-, (aR)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.



L4 ANSWER 4 OF 7 CA COPYRIGHT 2006 ACS on STN
 ACCESSION NUMBER: 142:373815 CA
 TITLE: Preparation of hydroxytrifluoromethylalkylpyrrolopyridines, -indoles, and related compounds as modulators of glucocorticoid receptor function

INVENTOR(S): Bekkali, Younes; Betageri, Rajashbar; Emmanuel, Michael J.; Hammach, Abdelhakim; Harcken, Hanke Justus Lee, Joechim; Kirrane, Thomas Martin; Kuzmich, Daniel; Thomas Wai-ho; Liu, Pingrong; Patel, Usha R.; Razavi, Hosseini; Riether, Doris; Takahashi, Hidenori; Thomson, David S.; Wang, Ji; Zindell, Renee; Proudfoot, John Robert Boehringer Ingelheim Pharmaceuticals, Inc., USA SOURCE: PCT Int. Appl. 549 pp. CODEN: PIIXD2 DOCUMENT TYPE: Patent LANGUAGE: English FAMILY ACC. NUM. COUNT: 1

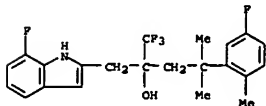
| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|---|------|----------|-----------------|------------|
| WO 2005030213 | A1 | 20050407 | WO 2004-US31009 | 20040923 |
| W: AS, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MO, MN, MM, MX, MZ, NA, NI, NO, NZ, OM, PO, PH, PL, PT, RO, RU, SC, SD, SE, SO, SK, SL, SY, TJ, TM, TN, TT, TZ, UA, UG, UZ, VC, VN, YU, ZA, ZM, ZW | | | | |
| RW: BM, GH, KE, LS, MM, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CO, CI, CM, GA, GN, GO, GW, ML, MR, NE, TD, TG | | | | |
| CA 2539909 | AA | 20050407 | CA 2004-2539909 | 20040923 |
| US 2005176706 | A1 | 20050811 | US 2004-947420 | 20040922 |
| PRIORITY APPLN. INFO.: | | | US 2003-505456P | P 20030924 |
| | | | US 2003-507079P | P 20030929 |
| | | | WO 2004-US31009 | W 20040922 |

OTHER SOURCE(S): MARPAT 142:373815
 AB Title compds., e.g. $\text{R}_1\text{R}_2\text{R}_3\text{OCH}_2\text{C}(\text{OH})\text{CP}_3\text{R}_4\text{R}_5$ (R_1 = (substituted) aryl, heteroaryl; R_2 , R_3 = H, alkyl; R_4R_5 atoms to form a C3=8 spiro cycloalkyl ring; R_4 = (substituted) alkyl, alkenyl, alkynyl; R_5 = substituted heteroaryl), were prepared for treatment of inflammatory, allergic, or proliferative processes (no date). Thus, N-[4-[6-(5-chloro-2,3-dihydrobenzofuran-7-yl)-4-hydroxy-6-methyl-4-trifluoromethylhept-1-ynyl]-2-isopropylpyrimidin-5-yl]-2,2,2-trifluoroacetamide (preparation given) and tetramethylguanidine were heated in

L4 ANSWER 4 OF 7 CA COPYRIGHT 2006 ACS on STN (Continued)
 dioxane at 100° for 1 h to give 4-(5-chloro-2,3-dihydrobenzofuran-7-yl)-1,1,1-trifluoro-2-(2-isopropyl-5H-pyrrolo[3,2-d]pyrimidin-6-ylmethyl)-4-methylpentan-2-ol.

IT 609850-91-1P
 RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PRSP (Preparation); USES (Uses)
 (preparation of hydroxytrifluoromethylalkylpyrrolopyridines, -indoles, and related compds. as modulators of glucocorticoid receptor function)

RN 609850-91-1 CA
 CN 1H-Indole-2-ethanol, 7-fluoro- α -(2-(5-fluoro-2-methylphenyl)-2-methylpropyl)- α -(trifluoromethyl)- (9CI) (CA INDEX NAME)



REFERENCE COUNT: 4 THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 5 OF 7 CA COPYRIGHT 2006 ACS on STN
 ACCESSION NUMBER: 141:140466 CA
 TITLE: Preparation of propenol and propylamine derivatives and their use as glucocorticoid ligands

INVENTOR(S): Proudfoot, John Robert; Regan, John Robinson; Thomson, David S.; Kuzmich, Daniel; Lee, Thomas Wai-ho; Hammach, Abdelhakim; Ralph, Mark Stephen; Zindell, Renee; Bekkali, Younes

PATENT ASSIGNEE(S): Boehringer Ingelheim Pharmaceuticals, Inc., USA SOURCE: PCT Int. Appl. 300 pp. CODEN: PIIXD2 DOCUMENT TYPE: Patent LANGUAGE: English FAMILY ACC. NUM. COUNT: 1

| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|---|------|----------|------------------|------------|
| WO 2004063163 | A1 | 20040729 | WO 2003-US40942 | 20031218 |
| W: AS, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MO, MN, MM, MX, MZ, NA, NI, NO, NZ, OM, PO, PH, PL, PT, RO, RU, SC, SD, SE, SO, SK, SL, SY, TJ, TM, TN, TT, TZ, UA, UG, UZ, VC, VN, YU, ZA, ZM, ZW | | | | |
| RW: BM, GH, GM, KE, LS, MM, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CO, CI, CM, GA, GN, GO, GW, ML, MR, NE, TD, TG | | | | |
| CA 2512257 | AA | 20040729 | CA 2003-2512257 | 20031218 |
| AU 2003297471 | A1 | 20040810 | AU 2003-297471 | 20031218 |
| US 2004162321 | A1 | 20040819 | US 2003-739208 | 20031218 |
| EP 1583745 | A1 | 20051012 | EP 2003-815237 | 20031218 |
| R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, SK | | | | |
| BR 2003017928 | A | 20051129 | BR 2003-17928 | 20031218 |
| CN 1735599 | A | 20060215 | CN 2003-80108207 | 20031218 |
| PRIORITY APPLN. INFO.: | | | US 2003-437925P | P 20030103 |
| | | | US 2003-445192P | P 20030205 |
| | | | WO 2003-US40942 | W 20031218 |

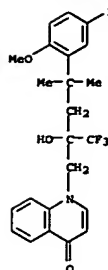
OTHER SOURCE(S): MARPAT 141:140466
 GI

AB Title compds. I [R1 = (heteroaryl), cycloalkyl, etc.; R2-3 = H, alkyl, arylalkyl, etc.; R4 = CO, divalent alkyl; R5 = pyrrolidine, morpholine, thiomorpholine, etc.; X = OH, amino] are prepared. For instance, 2-hydroxy-4-methyl-2-trifluoromethylpent-4-enic acid Et ester (preparation) is alkylated with 4-fluorobenzoic acid (AlCl₃); the resulting ester is reduced to the diol (LAH), converted to the oxirane (CH₂Cl₂, pyridine, NaCl) and treated with 2,6-dimethylmorpholine (DMF, 100°) to give II. I are glucocorticoid receptor modulators and are useful for the treatment of inflammatory disorders.

IT 727374-91-6, 1-[4-(5-fluoro-2-methoxyphenyl)-2-hydroxy-4-methyl-2-trifluoromethylpentyl]-1H-quinolin-4-one
RL: PAC (Pharmacological activity); RCT (Reactant); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); RACT (Reactant or reagent); USES (Uses)
 (preparation of propanol and propylamine derivs. and their use as glucocorticoid ligands)

RN 727374-91-6 CA

CN 4(1H)-Quinolinone, 1-[4-(5-fluoro-2-methoxyphenyl)-2-hydroxy-4-methyl-2-(trifluoromethyl)pentyl]- (9CI) (CA INDEX NAME)

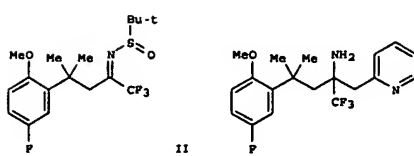
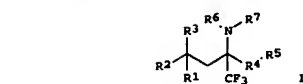


ACCESSION NUMBER: 140:27766 CA
TITLE: Preparation of 4-[(heteroaryl)-2-butylamine derivatives as glucocorticoid ligands
INVENTOR(S): Thomson, David; Kuzmich, Daniel; Kirrane, Thomas M.; Proudfoot, John Robert; Razavi, Hossein
PATENT ASSIGNEE(S): Boehringer Ingelheim Pharmaceuticals, Inc., USA
SOURCE: PCT Int. Appl. 122 pp.
CODEN: PIKXD2

DOCUMENT TYPE: Patent
LANGUAGE: English
FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION:

| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|---|------|----------|-----------------|-------------|
| WO 2003104195 | A1 | 20031218 | WO 2003-US17172 | 20030529 |
| W: AB, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KE, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MM, MX, MZ, NI, NO, NZ, OM, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, UZ, VC, VN, YU, ZA, ZM, ZW | | | | |
| RU: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, AM, AZ, BY, KG, KE, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CO, CI, CM, GA, GN, GU, GW, ML, MR, NE, SN, TD, TG | | | | |
| US 2004010148 | A1 | 20040115 | US 2003-446355 | 20030528 |
| CA 2486491 | AA | 20031218 | CA 2003-2486491 | 20030529 |
| AU 2003249669 | A1 | 20031222 | AU 2003-249669 | 20030529 |
| EP 1513810 | A1 | 20050316 | EP 2003-757304 | 20030529 |
| R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IB, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, ES, HU, SK | | | | |
| JP 2005529165 | T2 | 20050929 | JP 2004-511265 | 20030529 |
| US 2006014787 | A1 | 20060119 | US 2005-223501 | 20050909 |
| PRIORITY APPLN. INFO.: | | | US 2002-386334P | P 20020606 |
| | | | US 2003-446355 | A3 20030528 |
| | | | WO 2003-US17172 | W 20030529 |

OTHER SOURCE(S): MARPAT 140:27766
 GI



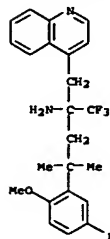
AB The title compds. I; R1 = (un)substituted (hetero)aryl; R2, R3 = H, alkyl; or R2 and R3 together with the carbon atom to which they are attached to form spiro cycloalkyl; R4 = alky, alkenyl, alkynyl; R5 = (un)substituted heteroaryl; R6, R7 = H, alkyl, alkenyl, alkoxy, etc., useful for modulating the glucocorticoid receptor function, and therefore for treating disease-states or conditions mediated by the glucocorticoid receptor function or characterized by inflammatory, allergic, or proliferative processes in a patient, were prepared and formulated.

Thus, treating 2-methylpyridine with tert-BuLi in THF followed by addition of the amide II (multi-step synthesis given) afforded 25% III which have shown activity as modulator of the glucocorticoid receptor function in one or more of the described in the patent assays (no specific data given). A kit for the in vitro diagnostic determination of the glucocorticoid receptor function is claimed.

IT 634203-47-7
RL: PAC (Pharmacological activity); RCT (Reactant); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); RACT (Reactant or reagent); USES (Uses)
 (preparation of 4-[(hetero)aryl]-2-butylamine derivs. as glucocorticoid ligands)

RN 634203-47-7 CA
CN 4-Quinolinetheanamine, α -(2-(5-fluoro-2-methoxyphenyl)-2-methylpropyl)- α -(trifluoromethyl)- (9CI) (CA INDEX NAME)

L4 ANSWER 6 OF 7 CA COPYRIGHT 2006 ACS on STN (Continued)



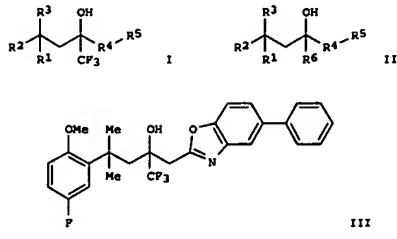
REFERENCE COUNT: 5 THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 7 OF 7 CA COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 139:292139 CA
TITLE: Preparation of heteroarylalkanols as glucocorticoid mimetics for treatment of inflammatory, allergic, and proliferative diseases
INVENTOR(S): Bekkali, Younes; Betageri, Raj; Gilmore, Thomas A.; Cardozo, Mario G.; Kirrane, Thomas M.; Kuzmich, Daniel; Proudfoot, John Robert; Takahashi, Hidenori; Thomson, David; Wang, Ji; Zindell, Renee; Harcken, Christian; Haeke Justus Joachim; Riehler, Doris
PATENT ASSIGNEE(S): Boehringer Ingelheim Pharmaceuticals, Inc., USA
SOURCE: PCT Int. Appl., 277 pp.
DOCUMENT TYPE: Patent
LANGUAGE: English
FAMILY ACC. NUM. COUNT: 1
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| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|---|------|----------|-----------------|-------------|
| WO 2003082280 | A1 | 20031009 | WO 2003-US8901 | 20030321 |
| W: AR, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, LZ, LC, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MM, MX, NZ, NI, NO, NZ, OM, PH, PL, PT, RO, RU, SC, SD, SE, SO, SK, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, UZ, VC, VN, YU, ZA, ZM, ZW | | | | |
| RU: GH, GM, KE, LS, KW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KO, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE, SI, SK, TR, SF, SJ, CF, CO, CM, GA, GH, GO, GW, ML, MR, NE, SN, TD, TO | | | | |
| CA 2478156 | AA | 20031009 | CA 2003-2478156 | 20030321 |
| AU 2003218342 | A1 | 20031013 | AU 2003-218342 | 20030321 |
| US 2004023999 | A1 | 20040205 | US 2003-394303 | 20030321 |
| US 6903215 | B2 | 20050607 | | |
| EP 1490062 | A1 | 20041229 | EP 2003-714339 | 20030321 |
| R: AT, BE, CH, DE, DK, ES, FR, GB, IR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, PI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, SK | | | | |
| BR 2003008784 | A | 20050111 | BR 2003-8784 | 20030321 |
| CN 1632296 | A | 20050629 | CN 2003-807180 | 20030321 |
| JP 2005527555 | T2 | 20050915 | JP 2003-579818 | 20030321 |
| US 2005059714 | A1 | 20050217 | US 2004-944615 | 20040917 |
| NO 200404031 | A1 | 20041019 | NO 2004-4031 | 20040924 |
| US 2005282881 | A1 | 20051222 | US 2005-185349 | 20050720 |
| PRIORITY APPLN. INFO.: US 2002-367758P | | | | P 20020326 |
| | | | US 2002-431817P | P 20021209 |
| | | | US 2003-442404P | P 20030124 |
| | | | US 2003-394303 | A1 20030321 |
| | | | WO 2003-US8901 | W 20030321 |
| | | | US 2004-944615 | A1 20040917 |

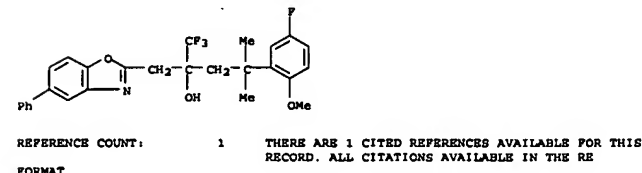
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OTHER SOURCE(S): MARPAT 139:292139

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AB Title compds. I and II [wherein R1 = substituted (hetero)aryl; R2 and R3 = independently H or alkyl; or CR2R3 = cycloalkyl; R4 = (un)substituted alkyl, alkenyl, or alkynyl; R5 = substituted heteroaryl; and R6 (when present) = (un)substituted alkyl, alkenyl, alkynyl, carbocyclic(alkyl), heterocyclic(alkyl), (hetero)aryl(alkyl), aryl(haloalkyl, carbocyclicalkenyl, heterocyclicalkenyl, or (hetero)arylalkenyl; and tautomers, prodrugs, solvates, or salts thereof] were prepared as glucocorticoid mimetics (no data). For example, 1,1,1-trifluoro-4-(5-fluoro-2-methoxyphenyl)-4-methylpentan-2-one (multi-step preparation from Et trifluoropyruvate, 1-bromo-2-methylpropene, and 4-fluoroanisole given) was coupled with 2-methyl-5-phenylbenzoxazole using LDA in THF to afford III. I, II, and pharmaceutical compns. containing such compns. are useful for treating inflammatory, allergic, or proliferative disorders mediated by glucocorticoid receptor (GR) function (no data).
IT 609849-72-1P, 1,1,1-Trifluoro-4-(5-fluoro-2-methoxyphenyl)-4-methyl-2-(5-phenylbenzoxazol-2-yl)methylpentan-2-ol
RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)
(glucocorticoid mimetic; preparation of heteroarylalkanols as GR modulators
for treatment of inflammatory, allergic, and proliferative diseases)
RN 609849-72-1 CA
CN 2-Benzoxazoleethanol, α -(2-(5-fluoro-2-methoxyphenyl)-2-methylpropyl)-5-phenyl- α -(trifluoromethyl)- (9CI) (CA INDEX NAME)

L4 ANSWER 7 OF 7 CA COPYRIGHT 2006 ACS on STN (Continued)



REFERENCE COUNT: 1 THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

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=> file caold

=> s l3
L5 0 L3

=> d his

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FILE 'REGISTRY' ENTERED AT 11:10:24 ON 06 JUN 2006
L1 STRUCTURE uploaded
L2 34 S L1 SAM
L3 629 S L1 FULL

FILE 'CA' ENTERED AT 11:10:51 ON 06 JUN 2006
L4 7 S L3

FILE 'CAOLD' ENTERED AT 11:11:04 ON 06 JUN 2006
L5 0 S L3

FILE 'MARPAT' ENTERED AT 11:11:11 ON 06 JUN 2006
L6 50 S L1

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STN INTERNATIONAL LOGOFF AT 11:14:55 ON 06 JUN 2006